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(54) REACTIVATION OF CARBON DIOXIDE ACCEPTORS

We, CONSOLIDATION COAL COM-PANY, a corporation organised under the laws of the State of Delaware, United States of America, of Koppers Building, Pittsburgh, 5 Pennsylvania, United States of America, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the

10 following statement: Carbon dioxide (CO2) is produced as an undesitable by-product in many processes. In some of these processes, it is desirable to remove the CO₂ as fast as it is formed. For that purpose, solids which readily react with CO₂ as the country of the CO₂ as fast as it.

CO2 may be used directly in the reaction zone, provided they do not react with the other reactants. Such solids are sometimes called "CO2 acceptors"

When lime (CaO) is used as the CO2 acceptor (and it is the favored acceptor because of its generally high effectiveness and low cost), the reaction of CaO and CO2 to form

CaCO, which is exothermic will provide the 25 requisite heat, or at least part of it, for endothermic reactions

A process wherein lime may be used to ad-vantage as an acceptor for CO2 involves the steam-carbon reaction which yields carbon 30 monoxide (CO), methane (CH₄) and hydrogen (H2), as well as the undesired CO2 as a by-product. Variants of the process are described in many patents.
United States Patent No. 2,705,672 is

35 illustrative. In this process, the heat generated by the reaction of CO₂ and CaO is nearly sufficient to supply that required for the endothermic reaction between the steam and carbon. As described in the cited patent, the 40 regeneration of the calcium oxide is accom-

regeneration of the carbonate to a tem-plished by heating the carbonate to a tem-perature of about 1700°F, if the pressure in the regenerator is atmospheric, or at or above 1900°F, if the pressure is at 20 atmospheres 45 or above. The calcium carbonate is thereby converted to CaO, and the CO2 evolved is

discharged for other purposes. The lime is recycled for re-use in the process.

Our experiments have clearly shown that the lime undergoes progressive decline in its ability to absorb CO₂ despite, or because of, its repeated cycling through the regeneration process. The decline in activity may be attri-buted to growth in crystallite size of the CaO, with consequent progressive reduction in pore volume on recycling through the steam-carbon process and associated regeneration step. In any case, the lime finally reaches the point where it is essentially inert so far as absorption of CO2 is concerned.

The primary object of the present invention is to provide a process of making an active CO2 acceptor from CaO which has become inert by frequent uses as above described, or which, in its natural state, is not

satisfactorily active. According to the invention, there is provided a process for regenerating active CaO, wherein CaO is used to absorb CO2 to form CaCO which is thereafter calcined to regenerate CaO for reuse in absorbing additional CO2, and wherein the CaO after repeated cycles of reuse and regeneration becomes deactivated or less effective in the absorption of CO2, which comprises converting said deactivated CaO to Ca(OH)₂, heating a mixture of said Ca(OH)₂ and either CaO or CaCO₃ or both in the presence of steam at an elevated pressure which is sufficiently high to prevent dehydration of the Ca(OH): until the mixture is melted, allowing the melted mixture to cool to a solid state, and calcining the solid to convert the Ca(OH)₂ to CaO, whereby an active CO. acceptor is formed.

In the various embodiments of the invention, the CaO is regenerated from the CaCO, by calcination at an elevated temperature, e.g. 1900°F. As pointed out above, the CaO is in an inactive state, the extent of inactivity being dependent upon the number of cycles of carbonation and regeneration it has been



subjected to.

The above process is based on the fact that active CaO may be obtained by reacting the inactive CaO obtained from CaCO,

with water and converting the Ca(OH)₂ thus obtained back to CaO. The overall process may be illustrated as follows:

$$CaCO_3 \rightarrow CO_2 + CaO$$
 (inactive form) $+ H_2O \rightarrow$
 $Ca(OH)_2 \rightarrow H_2O + CaO$ (active form)

Our invention which involves melting the mixture of Ca(OH)₂ and CaO and/or Ca(On, is also based upon the fact that Ca(OH)₂ forms low melting cutectic compositions with CaO, or CaCO₃, or both, as follows:

- 1) Ca(OH)₂.CaCO₃, melting point 1180°F. The phase diagram for this system is shown in Figure 1. 2) CaO.Ca(OH)₂, melting point—1454°F.
 - CaO.Ca(OH)₂, melting point—1454°F.
 The phase diagram for this system is shown in Figure 2.
- 6 3) CaO.Ca(OH)₂.CaCO₃.

In order to operate with compositions of the lowest possible melting points, it is obviously desirable to regulate the proportions of the components GO(BT), GAO, and GOO, to conform to those of the desired eutectics, although our process as outlined eutectics, although our process as outlined account in activity despite substantial deviation from the eutectic proportions.

30 In accordance with a preferred embodiment of this invention, a mixture of Ca(OH), and CaCO, in approximately 1:1 molar ratio is first obtained. This mixture may be obtained by any one of extend procedures including.

by any one of several procedures, including
the following:—

Procedure No. 1: Inactive CaO is converted
to Ca(OPL, by treatment with water (liquid,
not vapor) at relatively low temperature, cg.
300°F. This Ca(OFL), is then mixed with an
proximately equimolar amount of CaCO,
obtained from any source.

Procedure No. 2: Inactive CaO is converted to Ca(OH)₂ by treatment with steam in the presence of CaCO₂. The molar ratio, of Ca(OH)₂ to CaCO₂, if not 1:1, may be adjusted by adding CO₂.

The mixture of Ca(OH)s and CaCO, in 1:1 most ratio obtained by such procedures 50 is heated to a sufficiently high temperature in the presence of steam to a temperature from 1200°F; to 130°C. To melt the mixture. The 120°F; to 130°C. To melt the mixture. The prevent dehydration of the hydroxide and is generally 9—30 atmospheres.

The motion mixture is allowed to cool,

whereby a cutectic solid is formed as shown in Figure 1. The cutectic freezing point is 1180°F. To obtain the cutectic in a finely divided solid state, the molten mixture may simply be allowed to crystallize to a solid mass which is then ground and screened to recover the desired size; or the molten mixture cover the desired size; or the molten mixture

is allowed to drop through a cooling tower to from uniformly sized particles. The finely divided enterctic is subjected to low temperature dehydration at 1000 to 1100°F. to convert the Ca(OFD, to Ca(O, whereby a mixture of CaO and CaCO, is produced which may then be employed as the CO, acceptor. As assul, it has been found not only oppositely accept to the convertible of the convertible of

It is considered to be within the framework of the process of the present invention to blend warlows additives with the eutectic melt. Some additives such as various other acidic oxides, carbonates, phosphates, including by way of illustration, P.O., B.O., SiO., Al.O., Na,CO., may act as stabilizers to inhibit CaO crystal growth, Other additives may act as catalysts for the reactions in which the CO, acceptors are used.

acceptors are used.

A program was carried out to compare the activities of CaO compositions prepared from melts by the process of our invention with the activity of natural limestone.

The charge consisting of Ca(OFI), and Ca(O), was made up by hydrating Co(O) in liquid water and mixing with Ca(O), such that the proportion of Ca(OFI), to Ca(O), was known. The charge was heated in an autoclave to a temperature above the liquidus temperature, usually to about 1350°F. Excess steam pressure was vented.

The system was held at high temperature and pressure, usually 350 p.s.i.g, and 1350°F. For 20 minutes to ensure complete melting and was then cooled at the rate of 5°F. per minute to just below the eutectic temperature and then rapidly to room tem-

The frezen melt was then crushed and sized to particles of size going through a sieve of 14 × 48 mesh Tyler Standard and calcind to CaO in N, at 1600°P. in a fluidized bed reactor. The activity was then determined by recarbonation at 1 atmosphere in a fluidized bed at 1500°P. using pure CO. The activity copyresses as fractional curvation of GaO to CaCO, was determined control of the carbonate o

The activities of a number of melts prepared in this way from a natural limestone and from reagent grade materials are com-

pared in Table I below. It is seen that the activities of the melts, as long as they contain 40 mole per cent or more CaCO, are substantially identical to that obtained from the fresh limestone. A deactivated acceptor was prepared by cycling the fresh limestone 70 times through the calcining and gasification steps of the CO2 acceptor process de-

scribed in the previously cited U.S. Patent 2,705,672. The activity of the spent acceptor had dropped to 0.16. The inactive CO was converted to Ca(OH)₂ by treatment with hot water at 300°F, for one hour. The resulting mix was then melted and its activity determined by the procedure outlined above with results shown in Table I.

TABLE I Activity of Melts

Melt Composition Mole %

•	Title Composition Title /6			rictivity
Source of CaCO ₃	CaCO ₃	Ca(OH) ₂	CaO	After Melting
Fresh South Dakota Limestone	47	53		0.62
25 25 25 25	49	51		0.63
Deactivated Acceptor (Activity 0.16)	17 18	72 73	11 9	0.58 0.54
C.P. Reagent Chemicals	54 50 43 36 20	46 50 57 64 80		0.62 0.66 0.62 0.61 0.54

The activity of calcined fresh South Dakota Limestone is 0.66

It is seen that the acceptor was substantially. completely regenerated, i.e. its activity is 20 essentially indistinguishable from melts prepared from fresh materials of the same com-

The activity of CO₂ acceptors prepared by our process from South Dakota limestone was compared with that of the raw limestone by continuously recycling through a continuous bench scale unit for carrying our the CO₂ acceptor process described in U.S. Patent No. 2,705,672.

The acceptors were prepared by mixing Ca(OH)₂ with raw limestone. The mixture was heated to 1350°F, under 350 psig, steam pressure until it was completely melted. The frozen melt was crushed and sized to obtain a 16 × 28 mesh Tyler Standard acceptor for circulation through the continuous unit. The acceptor thus prepared contained 50 mole per cent Ca(OH)2 and 50 mole per cent CaCO2.

Prior to feeding the frozen melt to the continuous unit, the Ca(OH)2 was converted to CaO by heating at atmospheric pressure and 1000°F.

The raw limestone and the acceptors of this invention were recycled through the regenerator and gasifier vessels of the continuous unit under the conditions shown in Table II. The acceptors prepared by the melt process of this invention are referred to for convenience as melt acceptors.

TABLE II Conditions for Acceptor Activity Test

16 × 28 Mesh

Acceptor	South Dakota Limestone	16 × 28 Mesh Melt Acceptor		
Char	35 × 150 mesh devolatilized lignite char			
System Pressure Acceptor Circulation Rate (lb./hr.)	5.23	6.53		
Gasifier Conditions		•		
Temperature, °F. Char Feed Rate (lb./hr.) Char Bed Inventory, lb. Fluidizing Velocity, ft./sec.	7	00 6 8 27		
Outlet Gas Composition, Vol.	%			
H ₂ O H ₂ CH ₄ CO CO ₂ N ₂	1	8 5 0 6		
Regenerator Conditions Temperature, °F.	1910	1940		
Fluidizing Velocity, ft./sec. Outlet Gas Composition, Vol.	1. %	1		
CO ₂ N ₂	17.5 82.5	20 80		

The operation simulates in all details, a commercial CO2 acceptor system with the exception that the heat of calcination in the regenerator was supplied electrically instead of by in situ char combustion.

During the operation of the continuous unit, the acceptor enters the gasifier in the fully calcined condition and is recarbonated on of falling through the char bed. The driving force for recarbonation, i.e. Pco-Pco₂equil. is 0.5-0.8 atmospheres and the residence time of the acceptor is 7 minutes.

The activity is defined as the ratio moles CaCOs/total moles Ca in the acceptor leaving the gasifier.

The decline in activity of the limestone

and of the melt acceptor on recycling a number of times through the system for the two runs of Table II is shown in Figure 3. It is noted that the initial activities of the limestone and the melt acceptor containing 50 mole percent each of Ca(OH)₂ and CaCO₃ are substantially identical. The melt, however, declines more rapidly on recycling through the system. It should be noted, how-ever, in Table II that the regenerator temperature was about 30°F, higher in the melt acceptor run and this may be the cause of the more rapid decline in activity.

The higher temperature in the melt acceptor run was not accidental. The regenerator temperature was limited to 1910°F. in the limestone run to prevent agglomeration in the regenerator. The melt acceptor showed less tendency to agglomerate, which permitted an increase in regenerator temperature to 1940°F. The run also demonstrated one of the

other beneficial features of the melt acceptor, i.e. its high physical strength. The attrition rate on passing the acceptor through the pro-cess was extremely small, i.e. much less than

 weight per cent/cycle.
 A generalized type flowsheet which illustrates how the process may be conducted in practice is shown in Figure 4. The flowsheet is a process for reactivating an acceptor from

the steam-carbon reaction in which carbon monoxide, methane and hydrogen are formed and CO2 is a by-product and wherein CaO is used to absorb the CO2 produced and the resulting CaCO₃ is calcined to regenerate CaO
active CO₂ acceptor. The process is adapted
to produce a melt acceptor from either spent coarse acceptor, attrited acceptor fines, re-cycle fines from crushing and sizing of the melt, fresh limestone and any combination of

the above materials. The flowsheet as shown is illustrated for continuous operation, although the process may

tonination operation, annuaga are process may be conducted batchwise.

The feed in the feed hopper 10 will contain a mixture of CaO (from either spent acceptor or calcined limestone) and CaCOs. The feed mixture is fed to a melting unit 20 which is a pressurized stirred reactor to 20 which a mixture of steam and CO2 under pressure at 350—1000 psig. is also fed. The CaO is converted to Ca(OH)₂ by reaction with steam while a portion is converted to CaCOa by reaction with CO2. The relative 25 amounts of steam and CO2 reacted are adjusted to give the desired proportions of Ca(OH)₂ and CaCO₃ in the melt product, i.e. about 50 mole per cent of each.

The exothermic heats of the hydration and 30 carbonation reactions are sufficient to achieve the desired melting temperatures of 1200-1400°F., if the solid charge is preheated to

600-800°F

The melted acceptor is transferred to a prilling tower 30 through a sprayhead device 32. The molten spray is converted to a steady stream of droplets of approximately the proper size for use in the gasification plant, e. between 8 mesh and 28 mesh Tyler Standard. The melted acceptor is cooled and frozen by contact with a gas stream circulated under pressure through the prilling tower from a recirculation system 34. The Ca(OH)2 undergoes partial dehydration, and the endothermic heat of this reaction is sufficient to cause freezing of the melt,

An alternative arrangement is to spray the liquid melt into an immiscible liquid in which a temperature gradient is maintained such that the liquid drops freeze before passing out of the liquid. A possibility here, for example, is to spray the melt upward through a bath of molten lead

The frozen melt droplets are withdrawn to a hopper 40, and thence to a sieving and crushing system 50. The undersize material from crushing and sizing is returned to the melting unit. The oversize material is crushed.

The material of desired size is sent to a calcining unit 60 where the acceptor is calcined at 1000-1200°F, and atmospheric pressure to convert the Ca(OH)₂ to CaO be-

fore it is sent to the gasification plant.

WHAT WE CLAIM IS:-

1. In a process wherein CaO is used to absorb CO2 to form CaCO3 which is thereafter calcined to regenerate CaO for re-use in the process, and wherein the CaO after repeated re-use becomes deactivated or less effective in the absorption of CO2, the improvement for restoring said deactivated CaO to an effective CO₂ acceptor, which comprises the following steps: (a) converting said de-activated CaO to Ca(OH)₂, (b) heating a mix-ture of said Ca(OH)₂ and either CaO or CaCO₈ or both in the presence of steam at an elevated pressure which is sufficiently high to prevent dehydration of the Ca(OH)₂ until the mixture is melted, (c) allowing the melted mixture to cool to a solid state, and (d) calcining the solid from step (c) to convert the Ca(OH)₂ to CaO, whereby an active CO₂ ac-

ceptor is formed, 2. A process according to Claim 1 in which the inert CaO is converted in step (a) by treatment with liquid water. 3. A process according to Claim 1 in

which the mixture of step (b) is a mixture of Ca(OH)2 and CaCO3 in approximately 1:1

molar ratio. 4. A process according to Claims 1, 2 or 3 in which the CO2 is formed as a by-product when steam is reacted with carbon to

form CO, CH, and H2.

5. In the reaction of steam with carbon in which carbon monoxide, methane and hydrogen are formed and CO2 is a by-product and wherein CaO is used to absorb the CO2 produced and the resulting CaCO, is cal-cined to regenerate CaO active CO₂ acceptor, the improvement in the regeneration of the CaO active CO2-acceptor, which comprises the following steps: (a) introducing a mixture of deactivated CaO and CaCO, into a pressurized stirred reactor, (b) feeding a mixture of steam and CO2 to said stirred reactor while maintaining the pressure therein between 350 and 1000 psig and the temperature between 1200—1400°F, whereby part of the CaO is converted to Ca(OH)2 by reaction with the steam, and part is converted to CaCO, by reaction with CO2, the relative amounts of steam and CO2 reacted being amounts of sceam and was reacted using adjusted to yield a molten product containing about 50 mole percent of Ca(OH), and 50 mole percent of CaCO, (c) cooling the molten product from step (b) to a solid state, and (d) calcining the solid product to convert Ca(OH)₂ to CaO.

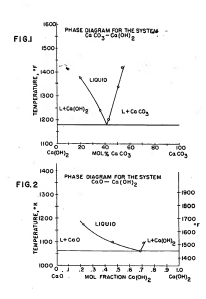
6. A process according to Claim 5 wherein the molten product from step (b) is passed through a prilling tower to form solid droplets.

7. A process for regenerating CaO in the active form as CO2-acceptor, from deactivated CaO, essentially as hereinbefore described and illustrated in the accompanying drawings.

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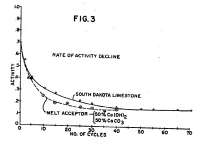
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